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(54) Recombinant immunoglobulin preparations

Rekombinante Immunoglobulin-Präparate
Préparations d'immunoglobuline recombinante

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[0001] This invintion relates to the field of immunoglobulin production and to modification of naturally occurring immunoglobulin amino acid sequences. Specifically, the invention relates to using ricombinant techniques to produce immunoglobulins which hav chimeric forms.

A. Immunoglobulinsand Antibodies

[0002] Antibodies are specific immunoglobulin polypeptides produced by the vertebrate immune system in response to challenge by foreign proteins, glycoproteins, cells, or other antigenic foreign substances. The sequence of events which permits the organism to overcome invasion by foreign cells or to rid the system of foreign substances is at least partially understood. An important part of this process is the manufacture of antibodies which bind specifically to a particular foreign substance. The binding specificity of such polypeptides to a particular antigen is highly refined, and the multitude of specificities capable of being generated by the individual vertebrate is remarkable in its complexity and variability. Thousands of antigens are capable of eliciting responses, each almost exclusively directed to the particular antigen which elicited it.

[0003] Immunoglobulins include both antibodies, as above described, and analogous protein substances which lack antigen specificity. The latter are produced at low levels by the lymph system and in increased levels by myelomas.

A.1 Source and Utility

[0004] Two major souces of vertebrate antibodies are presently utilized--generation in situ by the mammalian B lymphocytes and in cell culture by B-cell hybrids. Antibodies are made in situ as a result of the differentiation of immature B lymphocytes into plasma cells, which occurs in response to stimulation by specific antigens. In the undifferentiated B cell, the portions of DNA coding for the various regions on the immunoglobulin chains are separated in the genomic DNA. The sequences are reassembled sequentially prior to transcription. A review of this process has been given by Gough. Trends in Biochem Sci. 6: 203 (1981). The resulting rearranged genome is capable of expression in the mature B lymphocyte to produce the desired antibody. Even when only a single antigen is introduced into the sphere of the immune system for a particular mammal, however, a uniform population of antibodies does not result. The in situ immune response to any particular antigen is defined by the mosaic of responses to the various determinants which are present on the antigen. Each subset of homologous antibody is contributed by a single population of B cells--hence in situ generation of antibodies is "polyclonal".

[0005] This limited but inherent heterogeneity has been overcome in numerous particular cases by use of hybridoma technology to create "monoclonal" antibodies (Kohler, et al., Eur. J. Immunol., 6: 511 (1976)). In this process, splenocytes or lymphocytes from a mammal which has been injected with antigen are fused with a tumor cell line, thus producing hybrid cells or "hybridomas" which are both immortal and capable of producing the genetically coded antibody of the B cell. The hybrids thus formed are segregated into single genetic strains by selection, dilution, and regrowth. and each strain thus represents a single genetic line. They therefore produce immunoreactive antibodies against a desired antigen which are assured to be homogenous, and which antibodies, referencing their pure genetic parentage, are called "monoclonal". Hybridoma technology has to this time been focused largely on the fusion of munne lines, but humanhuman hybridomas (Olsson, L. et al., Proc. Natl. Acad. Sci. (USA). 77: 5429 (1980)): human-munne hybridomas (Schlom, J., et al. (ibid) 77: 6841 (1980)) and several other xenogenic hybrid combinations have been prepared as well. Alternatively, primary, antibody producing, B cells have been immortalized in vitro by transformation with viral DNA. [0006] Polyclonal, or, much more preferably, monoclonal, antibodies have a variety of useful properties similar to those of the present invention. For example, they can be used as specific immunoprecipitating reagents to detect the presence of the antigen which elicited the initial processing of the B cell genome by coupling this antigen-antibody reaction with suitable detection techniques such as labeling with radioisotopes or with enzymes capable of assay (RIA, EMIT, and ELISA). Antibodies are thus the foundation of immuno diagnostic tests for many antigenic substances. In another important use, antibodies can be directly injected into subjects suffering from an attack by a substance or organism containing the antigen in question to combat this attack. This process is currently in its experimental stages. but its potential is clearly seen. Third, whole body diagnosis and treatment is made possible because injected antibodies are directed to specific target disease tissues, and thus can be used either to determine the presence of the disease by carrying with them a suitable label, or to attack the diseased tissue by carrying a suitable drug.

[0007] Monoclonal antibodies produced by hybridomas, while theoretically effective as suggested above and clearly preferable to polyclonal antibodies because of their specificity, suffer from certain disadvantages. First, they tend to be contaminated with other proteins and cellular materials of hybridoma, (and, therefore, mammalian) origin. These cells contain additional materials, notably nucleic acid fragments, but protein fragments as well, which are capable of enhancing, causing, or mediating carcinogic responses. Second, hybridoma lines producing monoclonal antibodies

tend to be unstable and may alter the structure of antibody produced or stop producing antibody altogether (Kohler, G., et al., Proc. Natl. Acad. Sci (USA) 77: 2197 (1980); Morrison. S.L., J. Immunol. 123: 793 (1979)). The cell line genome appears to alter itself in respons to stimuli whose natur is not currently known, and this alteration may result in production of incorrect sequences. Third, both hybridoma and B cells inevitably produc certain antibodies in glycosylated form (M Ichers, F., Biochemistry, 10: 653 (1971)) which, under some circumstances, may be undesirable. Fourth, production of both monoclonal and polyclonal antibodies is relatively expensive. Fifth, and perhaps most important, production by current techniques (either by hybridoma or by B cell response) does not permit manipulation of the genome so as to produce antibodies with more effective design components than those normally elicited in response to antigens from the mature B cell in situ. The antibodies of the present invention do not suffer from the foregoing drawbacks, and, furthermore, offer the opportunity to provide molecules of superior design.

[0008] Even those immunoglobulins which lack the specificity of antibodies are useful, although over a smaller spectrum of potential uses than the antibodies themselves. In presently understood applications, such immunoglobulins are helpful in protein replacement therapy for globulin related anemia. In this context an inability to bind to antigen is in fact helpful, as the therapeutic value of these proteins would be impaired by such functionality. At present, such non-specific antibodies are derivable in quantity only from myeloma cell cultures suitably induced. The present invention offers an alternative, more economical source. It also offers the opportunity of cancelling out specificity by manipulating the four chains of the tetramer separately.

A.2 General Structure Characteristics

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[0009] The basic immunoglobin structural unit in vertebrate systems is now well understood (Edelman, G.M., Ann. N.Y. Acad. Sci., 190: 5 (1971)). The units are composed of two identical light polypeptide chains of molecular weight approximately 23,000 daltons, and two identical heavy chains of molecular weight 53.000-70,000. The four chains are joined by disulfide bonds in a "Y" configuration wherein the light chains bracket the heavy chains starting at the mouth of the Y and continuing through the divergent region as shown in figure 1. The "branch" portion. as there indicated, is designated the Tab region. Heavy chains are classified as gamma. mu, alpha, delta, or epsilon, with some subclasses among them, and the nature of this chain, as it has a long constant region, determines the "class" of the antibody as IgG, IgM, IgA, IgD, or IgE. Light chains are classified as either kappa or lambda. Each heavy chain class can be prepared with either kappa or lambda light chain. The light and heavy chains are covalently bonded to each other, and the "tail" portions of the two heavy chains are bonded to each other by covalent disulfide linkages when the immunoglobulins are generated either by hybridomas or by B cells. However, if non-covalent association of the chains can be effected in the correct geometry, the aggregate will still be capable of reaction with antigen, or of utility as a protein supplement as a non-specific immunoglobulin.

[0010] The amino acid sequence runs from the N-terminal end at the top of the Y to the C-terminal end at the bottom of each chain. At the N-terminal end is a variable region which is specific for the antigen which elicited it, and is approximately 100 amino acids in length, there being slight variations between light and heavy chain and from antibody to antibody. The variable region is linked in each chain to a constant region which extends the remaining length of the chain. Linkage is seen, at the genomic level, as occurring through a linking sequence known currently as the "J" region in the light chain gene, which encodes about 12 amino acids, and as a combination of "D" region and "J" region in the heavy chain gene, which together encode approximately 25 amino acids.

[0011] The remaining portions of the chain are referred to as constant regions and within a particular class do not to vary with the specificity of the antibody (i.e., the antigen eliciting it).

[0012] As stated above, there are five known major classes of constant regions which determine the class of the immunoglobulin molecule (IgG, IgM, IgA, IgD, and IgE corresponding γ , μ , α , δ , and ϵ heavy chain constant regions). The constant region or class determines subsequent effector function of the antibody, including activation of complement (Kabat, E.A., Structural Concepts in Immunology and Immunochemistry, 2nd Ed., p. 413-436, Holt, Rinehart, Winston (1976)), and other cellular responses (Andrews, D.W., et al., Clinical Immunobiology pp 1-18, W.B. Sanders (1980); Kohl. S., et al., Immunology. 48: 187 (1983)); while the variable region determines the antigen with which it will react.

B. Recombinant DNA Technology

[0013] Recombinant DNA technology has reached sufficient sophistication that it includes a repertoire of techniques for cloning and expression of gene sequenc s. Various DNA sequenc s can b r combined with some facility, creating new DNA entities capable of producing heterologous protein product in transform d microbes and cell cultures. The general means and methods for the <u>in vitro</u> ligation of various blunt ended or "sticky" ended fragments of DNA, for producing expression vectors, and for transforming organisms are now in hand.

[0014] DNA recombination of the essintial eliments (i.e., an origin of replication, on or mor phenotypic siliction

characteristics, expression control sequence, heterologous gene insert and remainder vector) generally is performed outside the host cell. The resulting recombinant replicable expression vector, or plasmid, is introduced into cells by transformation and large quantities of the recombinant vehicle is obtained by growing the transformant. Where the gene is properly inserted with reference to portions which govern the transcription and translation of the encoded DNA message, the resulting expression vector is useful to produce the polypeptide sequence for which the inserted gene codes, a process referred to as "expression." The resulting product may be obtained by lysis, if necessary, of the host cell and recovery of the product by appropriate purifications from other proteins.

[0015] In practice, the use of recombinant DNA technology can express entirely heterologous polypeptide-so-called direct expression--or alternatively may express a heterologous polypeptide fused to a portion of the amino acid sequence of a homologous polypeptide. In the latter cases, the intended bioactive product is sometimes rendered bioinactive within the fused, homologous heterologous polypeptide until it is cleaved in an extracellular environment

[0016] The art of maintaining cell or tissue cultures as well as microbial systems for studying genetics and cell physiology is well established. Means and methods are available for maintaining permanent cell lines, prepared by successive serial transfers from isolated cells. For use in research, such cell lines are maintained on a solid support in liquid medium, or by growth in suspension containing support nutriments. Scale-up for large preparations seems to pose only mechanical problems.

Summary of the Invention

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[0017] The present invention provides certain chimeric antibodies as set forth in the appended claim.

[0018] The invention can be used to prepare antibodies in pure "monoclonal" form. They can be manipulated at the genomic level to produce chimeras. They can also be manipulated at the protein level, since all four chains do not need to be produced by the same cell.

[0019] The invention is directed to. immunoglobulins which comprise polypeptides not hitherto found associated with each other in nature. "Fab proteins" may be produced which include only the "Fab" region of an immunoglobulin molecule i.e, the branches of the "Y". These Fab fragments may be chimeric, where for example, the constant and variable sequence patterns may be of different origin. Finally, either the light chain or heavy chain alone, or portions thereof, produced by recombinant techniques may be produced for the invention.

[0020] There are described expression vectors or plasmids capable of effecting the production of such immunoglobulins in suitable host cells. It includes the host cells and cell cultures which result from transformation with these vectors. Finally, there are described methods of producing these immunoglobulins and the DNA sequences, plasmids, and transformed cells intermediate to them.

Brief Description of the Drawings

[0021]

Figure 1 is a representation of the general structure of immunoglobulins.

Figure 2 shows the detailed sequence of the cDNA insert of pK17G4 which encodes kappa anti CEA chain.

Figure 3 shows the coding sequence of the fragment shown in Figure 2, along with the corresponding amino acid sequence.

Figure 4 shows the combined detailed sequence of the CDNA inserts of py298 and py11 which encode gamma anti CEA chain.

Figure 5 shows the corresponding amino acid sequence encoded by the fragment in Figure 4.

Figures 6 and 7 Outline the construction of expression vectors for kappa and gamma anti-CEA chains respectively. Figures 8A, 8B, and 8C show the results of sizing gels run on extracts of <u>E. coli</u> expressing the genes for gamma chain, kappa chain, and both kappa and gamma chains respectively.

Figure 9 shows the results of western blots of extracts of cells transformed as those in Figures 8.

Figure 10 shows a standard curve for ELISA assay of anti CEA activity.

Figures 11 and 12 show the construction of a plasmid for expression of the gene encoding a chimeric heavy chain. Figure 13 shows the construction of a plasmid for expression of the gene encoding the Fab region of heavy chain.

Detailed Description

A. Definitions

[0022] As us d her in, "antibodies" r fers to tetram rs or aggregates thereof which hav sp cific immunoreactiv activity, comprising light and heavy chains usually aggregat d in the "Y" configuration of Figur 1, with or without

covalent linkage between them; "immunoglobulins" refers to such assemblies whether or not specific immunareactive activity is a property. "Non-specific immunoglobulin" ("NSI") means those immunoglobulins which do not possess specificity-i. ., thos which are not antibodi s.

[0023] "Chimeric antibodi s" refers to those antibodies wherein one portion of each of the amino acid sequ nces of heavy and light chains is derived from a particular species, while the remaining segm int of the chains is derived from anoth r. Typically, in thes chimeric antibodi s, the variable region of both light and heavy chains is derived from antibodies from one species of mammals while the constant portions are derived from antibodies from another. One clear advantage to such chimeric forms is that, for example, the variable regions can conveniently be derived from presently known sources using readily available hybridomas or B cells from non human host organisms in combination with constant regions derived from, for example, human cell preparations. While the variable region has the advantage of ease of preparation, and the specificity is not affected by its source, the constant region being human, is less likely to elicit an immune response from a human subject when the antibodies are injected than would the constant region from a non-human source.

[0024] "Fab" region refers to those portions of the chains which are roughly equivalent, or analogous, to the sequences which comprise the Y branch portions of the heavy chain and to the light chain in its entirety, and which collectively (in aggregates) have been shown to exhibit antibody activity. "Fab protein", which protein is one of the aspects of the invention, includes aggregates of one heavy and one light chain (commonly known as Fab'), as well as tetramers which correspond to the two branch segments of the antibody Y, (commonly known as F(ab)₂), whether any of the above are covalently or non-covalently aggregated, so long as the aggregation is capable of selectively reacting with a particular antigen or antigen family. Fab antibodies have, been formed heretofore by proteolysis, and share the property of not eliciting antigen modulation on target tissues. However, as they lack the "effector" Fc portion they cannot effect, for example, lysis of the target cell by macrophages.

[0025] "Chimeric" Fab is defined analogously to the corresponding definition set forth in the previous paragraph for the Chimeric antibodies.

[0026] Individual heavy or light chains are "chimeric" in accordance with the above. As will become apparent from the detailed description of the invention, it is possible, using the techniques disclosed to prepare other combinations of the four-peptide chain aggregates, besides those specifically defined, such as hybrid antibodies containing chimeric light and mammalian heavy chains, hybrid Fab proteins containing chimeric Fab proteins of heavy chains associated with mammalian light chains, and so forth.

[0027] "Expression vector" includes vectors which are capable of expressing DNA sequences contained therein, i. e., the coding sequences are operably linked to other sequences capable of effecting their expression. It is implied, although not always explicitly stated, that these expression vectors must be replicable in the host organisms either as episomes or as an integral part of the chromosomal DNA. Clearly a lack of replicability would render them effectively inoperable. A useful, but not a necessary, element of an effective expression vector is a marker encoding sequence -- l.e. a sequence encoding a protein which results in a phenotypic property (e.g. tetracycline resistance) of the cells containing the protein which permits those cells to be readily identified. In sum, "expression vector" is given a functional definition, and any DNA sequence which is capable of effecting expression of a specified contained DNA code is included in this term, as it is applied to the specified sequence. As at present, such vectors are frequently in the form of plasmids, thus "plasmid" and "expression vector" are often used interchangeably. However, there are also described other forms of expression vectors which serve equivalent functions and which may, from time to time become known in the art

[0028] "Recombinant host cells" refers to cells which have been transformed with vectors constructed using recombinant DNA techniques. As defined herein, the antibody or modification thereof produced by a recombinant host cell is by virtue of this transformation, rather than in such lesser amounts, or more commonly, in such less than detectable amounts, as would be produced by the untransformed host

[0029] In descriptions of processes for isolation of antibodies from recombinant hosts, the terms "cell and "cell culture" are used interchangeably to denote the source of antibody unless it is clearly specified otherwise. In other words, recovery of antibody from the "cells" may mean either from spun down whole cells, or from the cell culture containing both the medium and the suspended cells.

B. Host Cell Cultures and Vectors

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[0030] The vectors and methods disclosed herein are suitable for use in host cells over a wide range of prokaryotic and ukaryotic organisms.

[0031] In general, of course, prokaryotes ar preferred for cloning of DNA's quenc's in constructing the vectors useful in the invention. For example, <u>E. coli</u> K12 strain 294 (ATCC No. 31446) is particularly useful. Other microbial strains which may be used include <u>E. coli</u> strains such as <u>E. coli</u> B, and <u>E. coli</u> X1776 (ATTC No. 31537). These examples are, of course, intended to be illustrative rather than limiting.

[0032] Prokaryotes may also be used for expression. The aforementioned strains, as well as $\underline{E.~coli}$ W3110 (F⁻, λ ⁻, prototrophic, ATTC No. 27325), bacilli such as $\underline{Bacillus~subtilus}$, and other enterobacteriaceae such as $\underline{Salmonella}$ typhimurium or S $\underline{rratia~marcesans}$, and various Pseudomonas species may be used.

[0033] In general, plasmid vectors containing r plicon and control sequences which are derived from species compatible with the host cell are used in connection with these hosts. The vector ordinarily carries a replication site, as well as marking sequences which are capable of providing phenotypic selection in transformed cells. For example, E. coli is typically transformed using pBR322, a plasmid derived from an E. coli species (Bolivar, et al., Gene 2: 95 (1977)). pBR322 contains genes for ampicillin and tetracycline resistance and thus provides easy means for identifying transformed cells. The pBR322 plasmid, or other microbial plasmid must also contain, or be modified to contain, promoters which can be used by the microbial organism for expression of its own proteins. Those promoters most commonly used in recombinant DNA construction include the β-lactamase (penicillinase) and lactose promoter systems (Chang et al. Nature, 275: 615 (1978); Itakura, et al, Science. 198: 1056 (1977); (Goeddel, et al Nature 281: 544 (1979)) and a tryptophan (trp) promoter system (Goeddel, et al, Nucleic Acids Res., 8: 4057 (1980); EPO Appl Publ No. 0036776). While these are the most commonly used, other microbial promoters have been discovered and utilized, and details concerning their nucleotide sequences have been published, enabling a skilled worker to ligate them functionally with plasmid vectors (Siebenlist, et al, Cell 20: 269 (1980)).

C. Methods Employed

C.1 Transformation:

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[0034] The preferred method of transfection is calcium treatment using calcium chloride as described by Cohen. F. N. et al. Proc. Natl. Acad. Sci. (USA), 69: 2110 (1972).

25 C.2 Vector Construction

[0035] Construction of suitable vectors containing the desired coding and control sequences employ standard ligation techniques. Isolated plasmids or DNA fragments are cleaved, tailored, and religated in the form desired to form the plasmids required. The methods employed are not dependent on the DNA source, or intended host.

[0036] Cleavage is performed by treating with restriction enzyme (or enyzmes) in suitable buffer. In general, about 1 μ g plasmid or DNA fragments is used with about 1 unit of enzyme in about 20 μ l of buffer solution. (Appropriate buffers and substrate amounts for particular restriction enzymes are specified by the manufacturer.) Incubation times of about 1 hour at 37° C are workable. After incubations, protein is removed by extraction with phenol and chloroform, and the nucleic acid is recovered from the aqueous fraction by precipitation with ethanol.

[0037] If blunt ends are required, the preparation is treated for 15 minutes at 15° with 10 units of <u>E. coli</u> DNA Polymerase I (Klenow), phenol-chloroform extracted, and ethanol precipitated.

[0038] Size separation of the cleaved fragments is performed using 6 percent polyacrylamide gel described by Goeddel, D., et al, <u>Nucleic Acids Res.</u>, 8: 4057 (1980) incorporated herein by reference.

[0039] For ligation, approximately equimolar amounts of the desired components, suitably end tailored to provide correct matching are treated with about 10 units T4 DNA ligase per 0.5 µg DNA. (When cleaved vectors are used as components, it may be useful to prevent religation of the cleaved vector by pretreatment with bacterial alkaline phosphatase.

[0040] In the examples described below correct ligations for plasmid construction are confirmed by transforming <u>E. coli</u> K12 strain 294 (ATCC 31446) with the ligation mixture. Successful transformants were selected by ampicillin or tetracycline resistance depending on the mode of plasmid construction. Plasmids from the transformants were then prepared, analyzed by restriction and/or sequenced by the method of Messing, et al, <u>Nucleic Acids Res.</u>, 9:309 (1981) or by the method of Maxam, et al, Methods in Enzymology, 65:499 (1980).

D. Outline of Procedures

D.1 Mammalian Antibodies

[0041] The first typ of antibody which is described her in is a "mammalian antibody"-one wherein the heavy and light chains are from an antibody oth rwis produced by a matur mammalian B lymphocyte eith r in situ or when fused with an immortalized c II as part of a hybridoma cultur. In outlin, the se antibodies are produced as follows:

[0042] Messenger RNA coding for heavy or light chain is isolated from a suitable source, either mature B cells or a hybridoma culture, employing standard techniques of RNA isolation, and the use of oligo-dT cellulose chromatography to s gregate the poly-A mRNA...The poly-A mRNA may, further, be fractionated to obtain sequences, of sufficient sizes.

to code for the amino acid sequences in the light or heavy chain of the desired antibody as the case may be.

[0043] A cDNA library is then pr par d from the mixture of mRNA using a suitable primer, preferably a nucleic acid sequ nce which is characteristic of the desired cDNA. Such a primer may be hypothesized and synthesized on the amino acid sequence of the antibody if the sequence is known. In the alternative cDNA from unfractionated poly-A mRNA from a cell line producing the desired antibody or poly-dT may also be used. The resulting cDNA is optionally size fractionated do no polyacrylamide gel and then extended with, for example, dC residues for annealing with pBR322 or other suitable cloning vector which has been cleaved by a suitable restriction enzyme, such as Pst I, and extended with dG residues. Alternative means of forming cloning vectors containing the cDNA using other tails and other cloning vector remainder may, of course, also be used but the foregoing is a standard and preferable choice. A suitable host cell strain, typically E. coli, is transformed with the annealed cloning vectors, and the successful transformants identified by means of, for example, tetracycline resistance or other phenotypic characteristic residing on the cloning vector plasmid.

[0044] Successful transformants are picked and transferred to microtiter dishes or other support for further growth and preservation. Nitrocellulose filter imprints of these growing cultures are then probed with suitable nucleotide sequences containing bases known to be complementary to desired sequences in the cDNA. Several types of probe may be used, preferably synthetic single stranded DNA sequences labeled by kinasing with ATP³². The cells fixed to the nitrocellulose filter are lysed, the DNA denatured, and then fixed before reaction with kinased probe. Clones which successfully hybridize are detected by contact with a photoplate, then plasmids from the growing colonies isolated and sequenced by means known in the art to verify that the desired portions of the gene are present.

[0045] The desired gene fragments are excised and tailored to assure appropriate reading frame with the control segments when inserted into suitable expression vectors. Typically, nucleotides are added to the 5' end to include a start signal and a suitably positioned restriction endonuclease site.

[0046] The tailored gene sequence is then positioned in a vector which contains a promoter in reading frame with the gene and compatible with the proposed host cell. A number of plasmids such as those described in U.S. Pat. Appln. Ser. Nos. 307473: 291892; and 305657 (EPO Publ. Nos. 0036776; 0048970 and 0051873) have been described which already contain the appropriate promoters, control sequences, ribosome binding sites, and transcription termination sites, as well as convenient markers.

[0047] The gene coding for the light chain and that coding for the heavy chain are recovered separately by the procedures outlined above. Thus they may be inserted into separate expression plasmids, or together in the same plasmid, so long as each is under suitable promoter and translation control.

[0048] The expression vectors constructed above are then used to transform suitable cells. The light and heavy chains may be transformed into separate cell cultures, either of the same or of differing species; separate plasmids for light and heavy chain may be used to co-transform a single cell culture, or, finally, a single expression plasmid containing both genes and capable of expressing the genes for both light and heavy chain may be transformed into a single cell culture.

[0049] Regardless of which of the three foregoing options is chosen, the cells are grown under conditions appropriate to the production of the desired protein. Such conditions are primarily mandated by the type of promoter and control systems used in the expression vector, rather than by the nature of the desired protein. The protein thus produced is then recovered from the cell culture by methods known in the art. but choice of which is necessarily dependent on the form in which the protein is expressed. For example, it is common for mature heterologous proteins expressed in \underline{E} . coli to be deposited within the cells as insoluble particles which require cell lysis and solubilization in denaturant to permit recovery. On the other hand, proteins under proper synthesis circumstances, in bacterial strains, can be secreted into the medium (gram positive bacteria) or into the periplasmic space (gram negative bactena) allowing recovery by less drastic procedures.

[0050] When heavy and light chain are coexpressed in the same host, the isolation procedure is designed so as to recover reconstituted antibody. This can be accomplished in vitro as described below, or might be possible in vivo in a microorganism which secretes the IgG chains out of the reducing environment of the cytoplasm. A more detailed description is given in D.2, below.

D.2 Chain Recombination Techniques

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[0051] The ability to produce heavy and light chains or portions thereof, in isolation from each other offers the opportunity to obtain unique and unprecedented assemblies of immunoglobulins and Fab regions. Such preparations require the use of techniques to reassemble isolated chains. Such means are known in the art, and it is, thus, appropriate to review them here.

[0052] While single chain disulfide bond containing prot ins have been r duced and reoxidized to regenerate in high yield native structure and activity (Freedman, R.B.. <u>t al</u>. In <u>Enzymology of Post Translational Modification of Proteins</u>, l: 157-212 (1980) Academic Press, NY.), proteins which consist of discontinuous polypeptide chains held togeth r by

disulfide bonds are more difficult to reconstruct in vitro after reductive cleavage. Insulin, a cameo case, has received much experimental attention over the years, and can now be reconstructed so efficiently that an industrial process has been built around it (Chance, R.E., et al., In Peptides: Proceedings of the Sev nth Annual American P ptide Symposium (Rich, D.H. and Gross, E., eds.) 721-728, Pierce Chemical Co., Rockford, IL. (1981)).

[0053] Immunoglobulin has proved a more difficult problem than insulin. The tetramer is stabilized intra and intermolecularly by 15 or more disulfide bonds. It has been possible to recombine heavy and light chains, disrupted by cleavage of only the interchain disulfides, to regain antibody activity even without restoration of the inter-chain disulfides (Edelman, G.M., et al., Proc. Natl. Acad. Sci. (USA) 50: 753 (1963)). In addition, active fragments of IgG formed by proteolysis (Fab fragments of 50.000 MW) can be split into their fully reduced heavy chain and light chain components and fairly efficiently reconstructed to give active antibody (Haber, E., Proc. Natl. Acad. Sci. (USA) 52: 1099 (1964); Whitney, P.L., et al., Proc. Natl. Acad. Sci. (USA) 53: 524 (1965)). Attempts to reconstitute active antibody from fully reduced native IgG have been largely unsuccessful, presumably due to insolubility of the reduced chains and of side products or intermediates in the refolding pathway (see discussion in Freedman, M.H., et al., J. Biol. Chem. 241: 5225 (1966)). If, however, the immunoglobulin is randomly modified by polyalanylation of its lysines before complete reduction, the separated chains have the ability to recover antigen-combining activity upon reoxidation (ibid).

[0054] A particularly suitable method for immunoglobulin reconstitution is derivable from the now classical insulin recombination studies, wherein starting material was prepared by oxidative sulfitolysis. thus generating thiol-labile S-sulfonate groups at all cysteines in the protein, non-reductively breaking disulfides (Chance et al. (supra)). Oxidative sulfitolysis is a mild disulfide cleavage reaction (Means, G.E.. et al., Chemical Modification of Proteins, Holden-Day, San Francisco (1971)) which is sometimes more gentle than reduction, and which generates derivatives which are stable until exposed to mild reducing agent at which time disulfide reformation can occur via thiol-disulfide interchange. The heavy and light chain S-sulfonates generated by oxidative sulfitolysis were reconstituted utilizing both air oxidation and thiol-disulfide interchange to drive disulfide bond formation. The general procedure is set forth in detail in U.S. Serial No. 452,187, filed Dec. 22, 1982 (EPO Appln. No. 83.307840.5). incorporated herein by reference.

D.3 Variants Permitted by Recombinant Technology

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[0055] Using the techniques described in paragraphs D.1 and D.2, additional operations which were utilized to gain efficient production of antibody can be varied in quite straightforward and simple ways to produce a great variety of modifications of this basic antibody form. These variations are inherent in the use of recombinant technology, which permits modification at a genetic level of amino acid sequences in normally encountered immunoglobulin chains, and the great power of this approach lies in its ability to achieve these variations, as well as in its potential for economic and specific production of desired scarce, and often contaminated, molecules. The variations also inhere in the ability to isolate production of individual chains, and thus create novel assemblies.

[0056] Briefly, since genetic manipulations permit reconstruction of genomic material in the process of construction of expression vectors, such reconstruction can be manipulated to produce new coding sequences for the components of "natural" antibodies or immunoglobulins. As discussed in further detail below, the coding sequence for a mammalian heavy chain may not be derived entirely from a single source or single species, but portions of a sequence can be recovered by the techniques described in D.1 from differing pools of mRNA, such as murine-murine hybridomas, human-murine hybridomas, or B cells differentiated in response to a series of antigen challenges. The desired portions of the sequences in each case can be recovered using the probe and analysis techniques described in D.1, and recombined in an expression vector using the same ligation procedures as would be employed for portions of the same model sequence. Such chimeric chains can be constructed of any desired length; hence, for example, a complete heavy chain can be constructed, or only sequence for the Fab region thereof.

[0057] The additional area of flexibility which arises from the use of recombinant techniques results from the power to produce heavy and light chains or fragments thereof in separate cultures or of unique combinations of heavy and light chain in the same culture, and to prevent reconstitution of the antibody or immunoglobulin aggregation until the suitable components are assembled. Thus, while normal antibody production results automatically in the formation of "mammalian antibodies" because the light and heavy chain portions are constructed in response to a particular determinant in the same cell, the methods of the present invention present the opportunity to assemble entirely new mixtures. Somewhat limited quantities of "hybrid" antibodies have been produced by "quadromas" i.e., fusions of two hybridoma cell cultures which permit random assemblies of the heavy and light chains so produced.

[0058] A more controlled ass mbly of desired chains is permitted, either by mixing the desired chains in vitro, or by transforming the sam culture with the coding sequences for the desired chains. ach other and mismatched to another pair gives the desired hybrid antibody.

D.4 Chimeric Antibodies

[0059] For construction of chimeric antibodies (wherein, for example, the variable's quences are separately derived from the constant sequences) the procedures of paragraph D.1 and D.2 are again applicable with appropriate additions and modifications. A preferred procedure is to recover desired portions of the genes encoding for parts of the heavy and light chains from suitable, differing, sources and then to religate these fragments using restriction endonucleases to reconstruct the gene coding for each chain.

[0060] For example, in a particularly preferred chimeric construction, portions of the heavy chain gene and of the light chain gene which encode the variable sequences of antibodies produced by a murine hybridoma culture are recovered and cloned from this culture and gene fragments encoding the constant regions of the heavy and light chains for human antibodies recovered and cloned from. for example, human myeloma cells. Suitable restriction enzymes may then be used to ligate the variable portions of the mouse gene to the constant regions of the human gene for each of the two chains. The chimeric chains are produced as set forth in D.1, aggregated as set forth in D.2 and used in the same manner as the non-chimeric forms. Of course, any splice point in the chains can be chosen.

D.5 Fab Protein

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[0061] Similarly, it is not necessary to include the entire gene for the heavy chain portion. All of the aforementioned variations can be superimposed on a procedure for Fab protein production and the overall procedure differs only in that that portion of the heavy chain coding for the amino terminal 220 amino acids is employed in the appropriate expression vector.

E. Specific Examples of Preferred Embodiments

[0062] The above description is in general terms and there follow several specific examples of embodiments which set forth details of experimental procedure in producing the desired antibodies. Example E.1 sets forth the general procedure for preparing anti CEA antibody components. Example E.3 sets forth the procedure for reconstitution and thus is applicable to preparation of chimeric immunoglobulins, and Fab proteins. Example E.4 sets forth the procedure for tailoring the heavy or light chain so that the variable and constant regions may be derived from different sources. Example E.5 sets forth the method of obtaining a shortened heavy chain genome which permits the production of the Fab regions and, in an analogous manner, Fc region.

E.1 Construction of Expression Vectors for Murine anti-CEA Antibody Chains and Peptide Synthesis

[0063] Carcinoembryonic antigen (CEA) is associated with the surface of certain tumor cells of human origin (Gold, 35 P., et al., J. Med., 122: 467 (1965)). Antibodies which bind to CEA (anti-CEA antibodies) are useful in early detection of these tumors (Van Nagell, T.R., et al., Cancer Res. 40: 502 (1980)), and have the potential for use in treatment of those human tumors which appear to support CEA at their surfaces. A mouse hybridoma cell line which secretes anti-CEA antibodies of the Igy1 class, CEA.66-E3, has been prepared as described by Wagener, C. et al., J. Immunol. 130, 2308 (1983) which is incorporated herein by reference, and was used as mRNA source. The production of anti 40 CEA antibodies by this cell line was determined. The N-terminal sequences of the antibodies produced by these cells was compared with those of monoclonal anti CEA as follows. Purified IgG was treated with PCAse (Podell, D.N., et al., Biochem. Biophys. Res. Commun. 81: 176 (1978)). and then dissociated in 6M guanidine hydrochloride, 10 mM 2-mercaptoethanol (1.0 mg of immunoglobulin. 5 min, 100° C water bath). The dissociated chains were separated on a Waters Associates alkyl phenyl column using a linear gradient from 100 percent A (0.1 percent TFA-water) to 90 percent B (TFA/H2O/MeCN 0.1/9.9/90) at a flow rate of 0.8 ml/min. Three major peaks were eluted and analyzed on SDS gels by silver staining. The first two peaks were pure light chain (Mu 25,000 daltons), the third peak showed a (7: 3) mixture of heavy and light chain. 1.2 nmoles of light chain were sequenced by the method of Shively, J.E., Methods in Enzymology, 79: 31 (1981), with an NH₂-terminal yield of 0.4 nmoles. A mixture of heavy and light chains (3 nmoles) was also sequenced, and sequence of light chain was deducted from the double sequence to yield the sequence of the heavy chain.

[0064] In the description which follows, isolation and expression of the genes for the heavy and light chains for anti CEA antibody produced by CEA.66-E3 are described. As the constant regions of these chains belong to the gamma and kappa families, respectively, "light chain" and "kappa chain", and "heavy chain" and "gamma chain", respectively, are used interchangeably below.

E.1.1 Isolation of Messenger RNA for Anti CEA Light and Heavy (Kappa and Gamma) Chains

[0065] Total RNA from CEA.66-E3 cells was xtract dessentially as report d by Lynch et al, Virology, 98: 251 (1979). C Ils wer pellet d by centrifugation and approximately 1 g portions of pellet r susp nded in 10 ml of 10 mM NaCl, 10 mM Tris HCl (pH 7.4), 1.5 mM MgCl₂. The resuspended cells were lysed by addition of non-ionic detergent NP-40 to a final concentration of 1 percent, and nuclei removed by centrifugation. After addition of SDS (pH 7.4) to 1 percent final concentration, the supernatant was extracted twice with 3 ml portions of phenol (redistilled)/chloroform: isoamyl alcohol 25:1 at 4°C. The aqueous phase was made 0.2 M in NaCl and total RNA was precipitated by addition of two volumes of 100 percent ethanol and overnight storage at -20°C. After centrifugation, polyA mRNA was purified from total RNA by oligo-dT cellulose chromatography as described by Aviv and Leder, Proc. Nat'l. Acad. Sci. (USA), 69: 1408 (1972). 142 µg of polyA mRNA was obtained from 1 g cells.

E.1.2 Preparation of E. coli Colony Library Containing Plasmids with Heavy and Light DNA Sequence Inserts

15 [0066] 5 μg of the unfractionated polyA mRNA prepared in paragraph E.1.1 was used as template for oligo-dT primed preparation of double-stranded (ds) cDNA by standard procedures as described by Goeddel et al., Nature 281: 544 (1979) and Wickens et al., J. Biol. Chem. 253: 2483 (1978) incorporated herein by reference. The cDNA was size fractionated by 6 percent polyacrylamide gel electrophoresis and 124 ng of ds cDNA greater than 600 base pairs in length was recovered by electroelution. A 20 ng portion of ds cDNA was extended with deoxy C residues using terminal deoxynucleotidyl transferase as described in Chang et al., Nature 275: 617 (1978) incorporated herein by reference, and annealed with 200 ng of the plasmid pBR322 (Bolivar et al., Gene 2: 95 (1977)) which had been cleaved with Pst I and tailed with deoxy G. Each annealed mixture was then transformed into E. coli K12 strain 294 (ATCC No. 31446). Approximately 8500 ampicillin sensitive, tetracycline resistant transformants were obtained.

25 E.1.3 Preparation of Synthetic Probes

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[0067] The 14mer, 5' GGTGGGAAGATGGA 3' complementary to the coding sequence of constant region for mouse MOPC21 kappa chain which begins 25 basepairs 3' of the variable region DNA sequence was used as kappa chain probe. A 15 mer, 5' GACCAGGCATCCCAG 3'. complementary to a coding sequence located 72 basepairs 3' of the variable region DNA sequence for mouse MOPC21 gamma chain was used to probe gamma chain gene. [0068] Both probes were synthesized by the phosphotriester method described in German Offenlegungschrift 2644432, incorporated herein by reference, and made radioactive by kinasing as follows: 250 ng of deoxyoligonucle-otide were combined in 25 μ l of 60 mM Tris HCI (pH 8), 10 mM MgCl₂. 15 mM betamercaptoethanol, and 100 μ Ci (γ -32P) ATP (Amersham, 5000 Ci.mMole). 5 units of T4 polynucleotide kinase were added and the reaction was allowed to proceed at 37°C for 30 minutes and terminated by addition of EDTA to 20 mM.

E.1.4 Screening of Colony Library for Kappa or Gamma Chain Sequences

[0069] ~2000 colonies prepared as described in paragraph E.1.2 were individually inoculated into wells of microtitre dishes containing LB (Miller, Experiments in Molecular Genetics, p. 431-3, Cold Spring Harbor Lab., Cold Spring Harbor, New York (1972)) + 5 μg, ml tetracycline and stored at -20°C after addition of DMSO to 7 percent. Individual colonies from this library were transferred to duplicate sets of Schfeicher and Schuell BA85/20 nitrocellulose filters and grown on agar plates containing LB + 5 µg ml tetracycline. After ~10 hours growth at 37°C the colony filters were transferred to agar plates containing LB + 5 μg ml tetracycline and 12.5 μg/ml chloramphenicol and reincubated overnight at 37°C. The DNA from each colony was then denatured and fixed to the filter by a modification of the Grunstein-Hogness procedure as described in Grunstein et al., Proc. Natl. Acad. Sci. (USA) 72: 3961 (1975). incorporated herein by reference. Each filter was floated for 3 minutes on 0.5 N NaOH, 1.5 M NaCl to lyse the colonies and denature the DNA then neutralized by floating for 15 minutes on 3 M NaCl. 0.5 M Tris HCl (pH 7.5). The filters were then floated for an additional 15 minutes on 2XSSC, and subsequently baked for 2 hours in an 80°C vacuum oven. The filters were prehybridized for ~2 hours at room temperature in 0 9 M NaCl. 1X Denhardts, 100 mM Tris HCl (pH 7.5), 5 mM Na-EDTA, 1 mM ATP, 1 M sodium phosphate (dibasic). 1 mM sodium pyrophosphate, 0.5 percent NP-40, and 200 μg/ ml E. coli t-RNA, and hybridized in the same solution overnight, essentially as described by Wallace et al. Nucleic Acids Research 9: 879 (1981) using -40×10⁶ cpm of either the kinased kappa or gamma probe described above. [0070] After ext_nsive washing at 37°C in 6X SSC. 0.1 percent SDS. th_filt_rs were_xpos_d to Kodak XR-5 X-ray film with DuPont Lightning-Plus intensifying screens for 124 hours at -80°C. Approximately 20 colonies which hybridized

with kappa chain probe and 20 which hybridized with gamma chain probe were characterized.

E.1.5 Characterization of Colonies which Hybridize to Kappa DNA Sequence Probe

[0071] Plasmid DNAs isolated from several different transformants which hybridiz d to kappa chain probe were cleaved with Pst I and fractionated by polyacrylamide gel electrophoresis (PAGE). This analysis d monstrat d that a number of plasmid DNAs contained cDNA inserts large enough to encode full length kappa chain. The complete nucleotide sequence of the cDNA insert of one of these plasmids was d t rmined by the dideoxynucleotide chain termination method as described by Smith. Methods Enzymol. 65, 560 (1980) incorporated herein by reference after subcloning restriction endonuclease cleavage fragments into M13 vectors (Messing et al., Nucleic Acids Research 9: 309 (1981). Figure 2 shows the nucleotide sequence of the cDNA insert of pK17G4 and Figure 3 shows the gene sequence with the corresponding amino acid sequence. Thus, the entire coding region of mouse anti-CEA kappa chain was isolated on this one large DNA fragment. The amino acid sequence of kappa chain, deduced from the nucleotide sequence of the pK17G4 cDNA insert, corresponds perfectly with the first 23 N-terminal amino acids of nature mouse anti-CEA kappa chain as determined by amino acid sequence analysis of purified mouse anti-CEA kappa chain. The coding region of pK17G4 contains 27 basepairs or 9 amino acids of the presequence and 642 basepairs or 214 amino acids of the mature protein. The mature unglycosylated protein (MW 24.553) has a variable region of 119 amino acids, including the J1 joining region of 12 amino acids, and a constant region of 107 amino acids. After the stop codon behind amino acid 215 begins 212 basepairs of 3' untranslated sequence up to the polyA addition. The kappa chain probe used to identify pK17G4 hybridizes to nucleotides 374-388 (figure 2).

20 E.1.6 Characterization of Colonies which Hybridize to Gamma 1 DNA Probe

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[0072] Plasmid DNA isolated from several transformants positive for hybridization with the heavy chain gamma 1 probe was subjected to Pst I restriction endonuclease analysis as described in E.1.5. Plasmid DNAs demonstrating the largest cDNA insert fragments were selected for further study. Nucleotide sequence coding for mouse heavy (gamma-1) chain, shows an Ncol restriction endonuclease cleavage site near the junction between variable and constant region. Selected plasmid DNAs were digested with both PstI and Ncol and sized on polyacrylamide. This analysis allowed identification of a number of plasmid DNAs that contain Ncol restriction endonuclease sites, although none that demonstrate cDNA insert fragments large enough to encode the entire coding region of mouse anti-CEA heavy chain.

[0073] In one plasmid isolated, p γ298 the cDNA insert of about 1300 bp contains sequence information for the 5' untranslated region, the signal sequence and the N-terminal portion of heavy chain. Because pγ298 did not encode the C-terminal sequence for mouse anti-CEA gamma 1 chain, plasmid DNA was isolated from other colonies and screened with Psti and Ncol. The C-terminal region of the cDNA insert of pγ11 was sequenced and shown to contain the stop codon, 3' untranslated sequence and that portion of the coding sequence missing from p γ298.

[0074] Figure 4 presents the entire nucleotide sequence of mouse anti-CEA heavy chain (as determined by the dideoxynucleotide chain termination method of Smith, Methods Enzymol., 65: 560 (1980)) and Figure 5 includes the translated sequence.

[0075] The amino acid sequence of gamma 1 (heavy chain) deduced from the nucleotide sequence of the $p\gamma298$ cDNA insert corresponds perfectly to the first 23 N-terminal amino acids of mature mouse anti-CEA gamma 1 chain as determined by amino acid sequence analysis of purified mouse anti-CEA gamma-1 chain. The coding region consists of 57 basepairs or 19 amino acids of presequences and 1346 basepairs or 447 amino acids of mature protein. The mature unglycosolated protein (MW 52.258) has a variable region of 135 amino acids, including a D region of 12 amino acids, and a J4 joining region of 13 amino acids. The constant region is 324 amino acids. After the stop codon behind amino acid 447 begins 96 bp of 3' untranslated sequences up to the polyA addition. The probe used to identify $P\gamma298$ and $P\gamma11$ hybridized to nucleotides 528-542 (Figure 4).

E.1.7 Construction of a Plasmid For Direct Expression of Mouse Mature Anti-CEA Kappa Chain Gene, pKCEAtrp207-1*

[0076] Figure 6 illustrates the construction of pKCEAtrp207-1*

[0077] First, an intermediate plasmid pHGH207-1*, having a single trp promoter, was prepared as follows:

[0078] The plasmid pHGH 207 (described in U.S. Pat Appl. Serial No. 307,473, filed Oct. 1, 1981 (EPO Publn. No. 0036776)) has a double lac promoter followed by the trp promoter, flanked by EcoR I sites and was used to prepare pHGH207-1. pHGH207 was digested with BamH 1, followed by partial digestion with EcoR I. The largest fragment, which contains the entire trp promoter, was isolated and ligated to the largest EcoR I- BamH I fragment from pBR322, and the ligation mixture used to transform E. coli 294. TetR AmpR colonies were isolated, and most of them contained pHGH207-1. pHGH207-1* which lacks the EcoR1 site between the ampR gene and the trp promoter, was obtained by partial digistion of pHGH207-1 with EcoR I, filling in the ends with Klenow and dNTPs, and religation.

Polymerase I in a 50 μ l reaction containing 60 mM NaCI, 7 mM MgCI₂, 7 mM Tris HCI (pH 7.4) and 1 mM in each dNTP at 37°C for 1 hour, followed by extraction with phenol/CHCI₃ and precipitation with ethanol. The precipitated DNA was digest d with BamH I, and the large vector fragment (fragment 1) purified using 5 perc nt polyacrylamid gel el ctrophoresis, electroelution, ph nol/CHCI₃ extraction and ethanol precipitation.

[0080] Th DNA was resuspended in 50 µl of 10 mM Tris pH 8, 1 mM EDTA and treated with 500 units Bacterial Alkaline Phosphatase (BAP) for 30' at 65° followed by phenol/CHCl₃ extraction and ethanol precipitation.

[0081] A DNA fragment containing part of the light chain sequence was prepared as follows: 7 µg of pK17G4 DNA was digested with Pst I and the kappa chain containing cDNA insert was isolated by 6 percent gel electrophoresis, and electroelution. After phenol/CHCl₃ extraction, ethanol precipitation and resuspension in water, this fragment was digested with Ava II. The 333 bp Pst I-Ava II DNA fragment was isolated and purified from a 6 percent polyacrylamide gel. [0082] A 15 nucleotide DNA primer was synthesized by the phosphotriester method G. O. 2.644.432 (supra) and has the following sequence: Met Asp IIe Val Met

5' ATG GAC ATT GTT ATG 3'

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[0083] The 5' methionine serves as the initiation codon. 500 ng of this primer was phosphorylated at the 5' end with 10 units T4 DNA kinase in 20 μ I reaction containing 0.5 mM ATP. ~200 ng of the Pst I-Ava II DNA fragment was mixed with the 20 μ I of the phosphorylated primer, heated to 95°C for 3 minutes and quick frozen in a dry-ice ethanol bath. The denatured DNA solution was made 60mM NaCI. 7mM MgCl₂. 7 mM Tris HCI (pH 7.4), 12 mM each dNTP and 12 units DNA Polymerase I-Large Fragment was added. After 2 hours incubation at 37°C this primer repair reaction was phenol. CHCl₃ extracted, ethanol precipitated. and digested to completion with Sau 3A. The reaction mixture was then electrophoresed on a 6 percent polyacrylamide gel and ~50 ng of the 182 basepair amino-terminal blunt-end to Sau 3A fragment (fragment 2) was obtained after electroelution.

[0084] 100 ng of fragment 1 (supra) and 50 ng of fragment 2 were combined in 20 μI of 20 mM Tris HCI (pH 7.5), 10 mM MgCl₂. 10 mM DTT, 2.5 mM ATP and 1 unit of T4 DNA ligase. After overnight ligation at 14°C the reaction was transformed into E. coli K12 strain 294, Restriction endonuclease digestion of plasmid DNA from a number of ampicillin resistant transformants indicated the proper construction and DNA sequence analysis proved the desired nucleotide sequence through the initiation codon of this new plasmid. pKCEAInt1 (Figure 6).

[0085] The remainder of the coding sequence of the kappa light chain gene was prepared as follows:

[0086] The Pst I cDNA insert fragment from 7 µg of K17G4 DNA was partially digested with Ava II and the Ava II cohesive ends were extended to blunt ends in a DNA Polymerase I large fragment reaction. Following 6 percent polyacrylamide gel electrophoresis the 686 basepair Pst I to blunt ended Ava II DNA fragment was isolated, purified and subjected to Hpa II restriction endonuclease digestion. The 497 basepair Hpa II to blunt ended Ava II DNA fragment (fragment 3) was isolated and purified after gel electrophoresis.

[0087] 10 μg of pKCEAInt1 DNA was digested with Ava I, extended with DNA polymerase I large fragment. and digested with Xba I. Both the large blunt ended Ava I to Xba I vector fragment and the small blunt ended Ava I to Xba I fragment were isolated and purified from a 6 percent polyacrylamide gel after electrophoresis. The large vector fragment (fragment 4) was treated with Bacterial Alkaline Phosphatase (BAP). and the small fragment was digested with Hpa II, electrophoresed on a 6 percent polyacrylamide and the 169 basepair Xba I-Hpa II DNA fragment (fragment 5) was purified. ~75 ng of fragment 4. ~50 ng of fragment 3 and ~50 ng of fragment 5 were combined in a T4 DNA ligase reaction and incubated overnight at 14°, and the reaction mixture transformed into E. coli K12 strain 294. Plasmid DNA from six ampicillin resistant transformants were analyzed by restriction endonuclease digestion. One plasmid DNA demonstrated the proper construction and was designated pKCEAInt2.

[0088] Final construction was effected by ligating the K-CEA fragment, including the trp promoter from pKCEAInt2 into pBR322(XAP). (pBR322(XAP) is prepared as described in U.S. Application 452.227. filed December 22, 1982; from pBR322 by deletion of the Aval-Pvull fragment followed by ligation.)

[0089] The K-CEA fragment was prepared by treating pKCEAInt2 with Ava I, blunt ending with DNA polymerase I (Klenow fragment) in the presence of DNTPs, digestion with Pst I and isolation of the desired fragment by gel electrophoresis and electroelution.

[0090] The large vector fragment from pBR322(XAP) was prepared by successive treatment with EcoR I. blunt ending with polymerase, and redigestion with Pst I, followed by isolation of the large vector fragment by electrophoresis and electroelution.

[0091] The K-CEA and large vector fragments as prepared in the preceding paragraphs were ligated with T4 DNA ligase, and the ligation mixture transformed into <u>E. coli</u> as above. Plasmid DNA from several ampicillin resistant transformants were sell cted for analysis, and one plasmid DNA demonstrated the proper construction, and was designated pKCEAtrp207-I*.

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E.1.8 Construction of a Plasmid Vector for Direct Expr ssion of Mouse Mature Anti-CEA Heavy (Gamma 1) Chain Gene, pyCEAtrp207-1*

[0092] Figur 7 Illustrates the construction of pγCEAtrp207-1*. This plasmid was constructed in two parts beginning with construction of the C-terminal region of the gamma 1 gene.

[0093] 5 μg of plasmid pHGH207-1* was digest d with Ava I, extended to blunt ends with DNA polym ras I large fragment (Klenow fragment), extracted with phenol CHCl₃. and ethanol precipitated. The DNA was digested with BamH I treated with BAP and the large fragment (fragment A) was purified by 6 percent polyacrylamide gel electrophoresis and electroelution.

[0094] ~5 μg of pγ11 was digested with Pst I and the gamma chain cDNA insert fragment containing the C-terminal portion of the gene was purified, digested with Ava II followed by extension of the Ava II cohesive ends with Klenow, followed by Taq I digestion. The 375 basepair blunt ended Ava II to Taq I fragment (fragment B) was isolated and purified by gel electrophoresis and electroelution.

[0095] 9 μg of pγ298 was digested with Tag I and BamH I for isolation of the 496 basepair fragment (fragment C).

[0096] Approximately equimolar amounts of fragments A, B, and C were ligated overnight at 14° in 20μl reaction mixture, then transformed into E. coli strain 294. The plasmid DNA from six ampicillin resistant transformants was committed to restriction endonuclease analysis and one plasmid DNA, named pγCEAInt, demonstrated the correct construction of the C-terminal portion of gamma 1 (Figure 5).

[0097] To obtain the N-terminal sequences, 30 μg of pγ298 was digested with Pst I and the 628 basepair DNA fragment encoding the N-terminal region of mouse anti-CEA gamma chain was isolated and purified. This fragment was further digested with Alu I and Rsa I for isolation of the 280 basepair fragment. A 15 nucleotide DNA primer met glu val met leu

5' ATG GAA GTG ATG CTG 3'

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was synthesized by the phosphotriester method (supra).

[0098] The 5' methionine serves as the initiation codon. 500 ng of this synthetic oligomer primer was phosphorylated at the 5' end in a reaction with 10 units T4 DNA kinase containing 0.5 mM ATP in 20μl reaction mixture. ~500 ng of the 280 basepair Alu I-Rsa I DNA fragment was mixed with the phosphorylated primer. The mixture was heat denatured for 3 minutes at 95° and quenched in dry-ice ethanol. The denatured DNA solution was made 60mM NaCl, 7mM MgCl₂, 7 mM Tris HCl (pH 7.4), 12 mM in each dNTP and 12 units DNA Polymerase I-Large Fragment was added. After 2 hours incubation at 37°C, this primer repair reaction was phenol/CHCl₃ extracted, ethanol precipitated, and digested to completion with Hpall. ~50 ng of the expected 125 basepair blunt-end to HPa II DNA fragment (fragment D) was purified from the gel.

[0099] A second aliquot of py298 DNA was digested with Pst I, the 628 basepair DNA fragment purified by polyacrylamide gel electrophoresis, and further digested with BamH I and Hpa II. The resulting 380 basepair fragment (fragment E) was purified by gel electrophoresis.

[0100] -5 μg of pγCEAntl was digested with EcoR I, the cohesive ends were made flush with DNA polymerase I (Klenow), further digested with BamH I, treated with BAP and electrophoresed on a 6 percent polyacrylamide gel. The large vector fragment (fragment F) was isolated and purified.

[0101] In a three fragment ligation, 50 ng fragment D, 100 ng fragment E, and 100 ng fragment F were ligated overnight at 4° in a 20 μ l reaction mixture and used to transform E. coli K12 strain 294. The plasmid DNAs from 12 ampicillin resistant transformants were analyzed for the correct construction and the nucleotide sequence surrounding the initiation codon was verified to be correct for the plasmid named pyCEAInt2.

[0102] The expression plasmid, $p\gamma$ CEAtrp207-I* used for expression of the heavy chain gene is prepared by a 3-way ligation using the large vector fragment from pBR322(XAP) (supra) and two fragments prepared from p γ CEAInt2.

[0103] pBR322(XAP) was treated as above by digestion with EcoR1, blunt ending with DNA polymerase (Klenow) in the presence of dNTPs, followed by digestion with Pst I, and isolation of the large vector fragment by gel electrophoresis. A 1543 base pair fragment from pγCEAInt2 containing trp promoter linked with the N-terminal coding region of the heavy chain gene was isolated by treating pγCEAInt2 with Pst I followed by BamH I, and isolation of the desired fragment using PAGE. The 869 base pair fragment containing the C-terminal coding portion of the gene was prepared by partial digestion of pγCEAInt2 with Ava I, blunt ending with Klenow, and subsequent digestion with BamH I, followed by purification of the desired fragment by gel electrophoresis.

[0104] The aforementioned three fragments were then ligated under standard conditions using T4 DNA ligase, and a ligation mixture used to transform \underline{E} . \underline{coli} strain 294. Plasmid DNAs from several tetracycline resistant transformants were analyzed; on plasmid DNA demonstrated the proper construction and was designat d pyCEAtrp207-1*.

E.1.9 Production of Immunoglobulin Chains by E. coli

[0105] E. coli strain W3110 (ATTC No. 27325) was transformed with pyCEAtrp207-1* or pKCEAtrp207-1* using stand-

ard techniques.

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[0106] To obtain double transformants, <u>E. coli</u> strain W3110 cells were transformed with a modified pKCEAtrp207-1*, pKCEAtrp207-1*Δ, which had been modified by cleaving a Pst I-Pvu I fragm int from the amp ^R gene and religating. Cells transformed with pKCEAtrp207-1*Δ are thus sensitive to ampicillin but still resistant to tetracycline. Successful transformants were retransformed using pγCEAInt2 which confers resistance to ampicillin but not tetracycline. Cells containing both pKCEAtrp207-1*Δ and pγCEAInt2 thus identified by growth in a medium containing both ampicillin and tetracycline.

[0107] To confirm the production of heavy and/or light chains in the transformed cells, the cell samples were inoculated into M9 tryptophan free medium containing 10μg ml tetracycline, and induced with indoleacrylic acid (IAA) when the OD 550 reads 0.5. The induced cells were grown at 37°C during various time periods and then spun down, and suspended in TE buffer containing 2 percent SDS and 0.1 M β-mercaptoethanol and boiled for 5 minutes. A 10 x volume of acetone was added and the cells kept at 22°C for 10 minutes, then centrifuged at 12,000 rpm. The precipitate was suspended in O'Farrell SDS sample buffer (O'Farrell, P.H., J. Biol. Chem., 250: 4007 (1975)); boiled 3 minutes, recentrifuged, and fractionated using SDS PAGE (10 percent), and stained with silver stain (Goldman, D. et al., Science 211: 1437 (1981)): or subjected to Western blot using rabbit anti-mouse IgG (Burnett, W. N., et al., Anal. Biochem. 112: 195 (1981)). for identification light chain and heavy chain.

[0108] Cells transformed with pyCEAtrp207-1* showed bands upon SDS PAGE corresponding to heavy chain molecular weight as developed by silver stain. Cells transformed with pKCEAtrp207-1* showed the proper molecular weight band for light chain as identified by Western Blot: double transformed cells showed bands for both heavy and light chain molecular weight proteins when developed using rabbit anti-mouse IgG by Western blot. These results are shown in Figures 8A, 8B, and 8C.

[0109] Figure 8A shows results developed by silver stain from cells transformed with pγCEAtrp207-1*. Lane 1 is monoclonal anti-CEA heavy chain (standard) from CEA.66-E3. Lanes 2b-5b are timed samples 2 hrs. 4 hrs, 6 hrs, and 24 hrs after IAA addition. Lanes 2a-5a are corresponding untransformed controls: Lanes 2c-5c are corresponding uninduced transformants.

[0110] Figure 8B shows results developed by Western blot from cells transformed with pKCEAtrp207-1*. Lanes 1b-6b are extracts-from induced cells immediately, 1hr, 3.5 hrs. 5 hrs. 8 hrs. and 24 hrs after IAA addition. and 1a-6a corresponding uninduced controls. Lane 7 is an extract from a pγCEAtrp207-1* control, lanes 8. 9, and 10 are varying amounts of anti CEA-kappa chain from CEA.66-E3 cells.

[0111] Figure 8C shows results developed by Western blot from four colonies of double transformed cells 24 hours after IAA addition (lanes 4-7]. Lanes 1-3 are varying amounts of monoclonal gamma chain controls, lanes 6 and 9 are untransformed and pyCEAtrp207-1* transformed cell extracts, respectively.

[0112] In another quantitative assay, frozen, transformed <u>E. coli</u> cells grown according to E.1.10 (below) were lysed by heating in sodium dodecyl sulfate (SDS)/β-mercaptoethanol cell lysis buffer at 100°. Aliquots were loaded on an SDS polyacrylamide gel next to lanes loaded with various amounts of hybndoma anti-CEA. The gel was developed by the Western blot. Burnett (supra), using ¹²⁵I-labeled sheep anti-mouse IgG antibody from New England Nuclear. The results are shown in Figure 9. The figure shows that the <u>E. coli</u> products co-migrate with the authentic hybridoma chains, indicating no detectable proteolytic degradation in <u>E. coli</u>. Heavy chain from mammalian cells is expected to be slightly heavier than <u>E. coli</u> material due to glycosylation in the former. Using the hybridoma lanes as a standard, the following estimates of heavy and light chain production were made:

	(Per gram of cells)
E. coli (W3110/pγCE Atr p207-1*)	5 mg γ
E. coli (W3110/pKCE Atr p207-1*)	1.5 mg K
E. coli (W3110/pKCEAtrp207-1*Δ, pγCEAInt2)	0.5 mg K, 1.0 mg γ

E.1.10 Reconstitution of Antibody from Recombinant K and Gamma Chains

[0113] In order to obtain heavy and light chain preparations for reconstitution, transformed cells were grown in larger batches, harvested and frozen. Conditions of growth of the variously transformed cells were as follows:

[0114] <u>E. coli</u> (W3110/ργCEAtrp207-1* were inoculated into 500 ml LB medium containing 5μg ml tetracycline and grown on a rotary shaker for 8 hours. The culture was then transferred to 10 liters of fermentation medium containing yeast nutrients, salts, glucose, and 2μg/ml tetracyclin. Additional glucose was add d during growth aid at OD 550 = 20, indoleacrylic (IAA), a trp derepressor, was added to a concentration of 50 μg/ml. The cills wer if d additional glucose to a final OD 550 = 40, achieved approximating ly 6 hours from the IAA addition.

[0115] E. coli (W3110) cells transform d with pKCEA trp 207-1* and double transform d (with pKCEAtrp207-1* Δ and pyCEAInt2) were grown in a manner analogous to that described above except that the OD 550 six hours after

IAA addition at harvest was 25-30.

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[0116] The cells were then harv st d by centrifugation, and frozen.

E.2 Assay Method for Reconstitut d Antibody

[0117] Anti-CEA activity was d termin d by ELISA as a crit rion for succ ssful r constitution. Wells of microtit r plates (Dynatech Immulon) were saturated with CEA by incubating 100 µl of 2-5 µg CEA/ml solution in 0.1 M carbonate buffer, pH 9.3 for 12 hours at room temperature. The wells were then washed 4 times with phosphate buffered saline (PBS), and then saturated with BSA by incubating 200 µl of 0.5 percent BSA in PBS for 2 hours at 37°C, followed by washing 4 times with PBS. Fifty microliters of each sample was applied to each well. A standard curve (shorn in Figure 10), was run, which consisted of 50 µl samples of 10 µg, 5 µg, 1 µg, 500 ng, 100 ng, 50 ng, 10 ng, 5 ng and 1 ng anti-CEA/ml in 0.5 percent BSA in PBS, plus 50 µl of 0.5 percent BSA in PBS alone as a blank. All of the samples were incubated in the plate for 90 minutes at 37°C.

[0118] The plates were then washed 4 times with PBS, and sheep anti-mouse IgG-alkaline phosphate (TAGO, Inc.) was applied to each well by adding 100 μ I of an enzyme concentration of 24 units/mI in 0.5 percent BSA in PBS. The solution was incubated at 37°C for 90 minutes. The plates were washed 4 times with PBS before adding the substrate. 100 μ I of a 0.4 mg/mI solution of p-nitrophenylphosphate (Sigma) in ethanolamine buffered saline, pH 9.5. The substrate was incubated 90 minutes at 37°C for color development.

[0119] The A_{450} of each well was read by the Microelisa Auto Reader (Dynatech) set to a threshold of 1.5, calibration of 1.0 and the 0.5 percent BSA in PBS (Blank) well set to 0.000. The A_{450} data was tabulated in RS-1 on the VAX system, and the standard curve data fitted to a four-parameter logistic model. The unknown samples concentrations were calculated based on the A_{450} data.

E.3 Reconstitution of Recombinant Antibody and Assay

[0120] Frozen cells prepared as described in paragraph E.1.10 were thawed in cold lysis buffer [10mM Tris HCI, pH 7.5, 1mM EDTA, 0.1 M NaCI, 1mM phenylmethylsulfonyl fluoride (PMSF)] and lysed by sonication. The lysate was partially clarified by centrifugation for 20 mins at 3,000 rpm. The supernatant was protected from proteolytic enzymes by an additional 1 mM PMSF, and used immediately or stored frozen at -80°C; frozen lysates were never thawed more than once.

[0121] The S-sulfonate of <u>E. coli</u> produced anti-CEA heavy chain (γ) was prepared as follows: Recombinant <u>E. coli</u> cells transformed with pγCEAtrp207-1* which contained heavy chain as insoluble bodies, were lysed and centrifuged as above; the pellet was resuspended in the same buffer, sonicated and re-centrifuged. This pellet was washed once with buffer, then suspended in 6M guanidine HCI, 0.1M Tris HCI, pH 8, 1mM EDTA, 20 mg/ml sodium sulfite and 10 mg/ml sodium tetrathionate and allowed to react at 25° for about 16 hrs. The reaction mixture was dialyzed against 8M urea, 0.1 M Tris HCI, pH 8, and stored at 4°, to give a 3 mg/ml solution of _γ-SSO₃.

[0122] 650 μl of cell lysate from cells of various $\underline{E.~coli}$ strains producing various lgG chains, was added to 500 mg urea. To this was added β -mercaptoethanol to 20mM, Tris-HCI, pH 8.5 to 50mM and EDTA to 1mM, and in some experiments, γ -SSO₃ was added to 0.1 mg/ml. After standing at 25° for 30-90 mins., the reaction mixtures were dialyzed at 4° against a buffer composed of 0.1M sodium glycinate, pH 10.8, 0.5M urea, 10mM glycine ethyl ester, 5mM reduced glutathione, 0.1 mM oxidized glutathione. This buffer was prepared from N₂-saturated water and the dialysis was performed in a capped Wheaton bottle. After 16-48 hours, dialysis bags were transferred to 4° phosphate buffered saline containing 1mM PMSF and dialysis continued another 16-24 hrs. Dialysates were assayed by ELISA as described in paragraph E.2 for ability to bind CEA. The results below show the values obtained by comparison with the standard curve in x ng/ml anti-CEA. Also shown are the reconstitution efficiencies calculated from the ELISA responses, minus the background (108 ng/ml) of cells producing K chain only, and from estimates of the levels of γ and K chains in the reaction mixtures.

	ng/ml anti-CEA	Percent recombination
E. coli W3110 producing IFN-αA (control)	0	
E. coli (M3110/pKCEAtrp207-1*).	108	
E. coli (M3110/pKCEAtrp207-1*), plus γ-SSO ₃	848	0.33
E. coli (W3110/pKCEAtrp207-1*Δ, pγCEAInt2)	1580	0.76
Hybridoma anti-CEA K-SSO ₃ and γ-SSO ₃	540	0.40

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E.4 Preparation of Chimeric Antibody

[0123] Figur s 11 and 12 show the construction of an expression vector for a chimeric heavy (gamma) chain which comprises the murine anti CEA variable region and human γ -2 constant region.

- [0124] A DNA sequence encoding the human gamma-2 heavy chain is prepared as follows: the cDNA library obtain d by standard techniques from a human multiple myeloma cell line is probed with 5' GGGCACTC-GACACAA 3' to obtain the plasmid containing the cDNA insert for human gamma-2 chain (Takahashi, et al., Cell, 29: 671 (1982), incorporated herein by reference), and analyzed to verify its identity with the known sequence in human gamma-2 (Ellison, J., et al., Proc. Natl. Acad. Sci. (USA), 79: 1984 (1982) incorporated herein by reference).
- [0125] As shown in Figure 11, two fragments are obtained from this cloned human gamma 2 plasmid (ργ2). The first fragment is formed by digestion with Pvuli followed by digestion with Ava III, and purification of the smaller DNA fragment, which contains a portion of the constant region, using 6 percent PAGE. The second fragment is obtained by digesting the pγ2 with any restriction enzyme which cleaves in the 3' untranslated region of γ2, as deduced from the nucleotide sequence, filling in with Klenow and dNTPs, cleaving with Ava III, and isolating the smaller fragment using 6 percent PAGE. (The choice of a two step, two fragment composition to supply the Pvull-3' untranslated fragment provides a cleaner path to product due to the proximity of the AvallI site to the 3 terminal end thus avoiding additional restriction sites in the gene sequence matching the 3' untranslated region site.) pγCEA207-1* is digested with EcoR 1, treated with Klenow and dNTPs to fill in the cohesive end, and digested with Pvu II, the large vector fragment containing promoter isolated by 6 percent PAGE.
- [0126] The location and DNA sequence surrounding the Pvull site in the mouse gamma-1 gene are identical to the location and DNA sequence surrounding the Pvull site in the human gamma-2 gene.
 - [0127] The plasmid resulting from a three way ligation of the foregoing fragments, pChim1. contains. under the influence of trp promoter, the variable and part of the constant region of murine anti-CEA gamma 1 chain. and a portion of the gamma 2 human chain. pChim1 will. in fact. express a chimeric heavy chain when transformed into E. coli, but one wherein the change from mouse to human does not take place at the variable to constant junction.
 - [0128] Figure 12 shows modification or pChim1 to construct pChim2 so that the resulting protein from expression will contain variable region from murine anti CEA antibody and constant region from the human γ -2 chain. First, a fragment is prepared from pChim1 by treating with Nco I, blunt ending with Klenow and dNTPs, cleaving with Pvu II, and isolating the large vector fragment which is almost the complete plasmid except for short segment in the constant coding region for mouse anti CEA. A second fragment is prepared from the previously described py2 by treating with Pvu II, followed by treating with any restriction enzyme which cleaves in the variable region, blunt ending with Klenow and dNTPs and isolating the short fragment which comprises the junction between variable and constant regions of this chain.
 - [0129] Ligation of the foregoing two fragments produces an intermediate plasmid which is correct except for an extraneous DNA fragment which contains a small portion of the constant region of the munne anti CEA antigen, and a small portion of the variable region of the human gamma chain. This repair can be made by excising the Xba I to Pvu II fragment and cloning into M13 phage as described by Messing et al., Nucleic Acids Res. 9: 309 (1981), followed by in vitro site directed deletion mutagenesis as described by Adelman, et al., DNA 2, 183 (1983) which is incorporated herein by reference. The Xba I-Pvu II fragment thus modified is ligated back into the intermediate plasmid to form pChim2. This plasmid then is capable of expressing in a suitable host a cleanly constructed murine variable human constant chimeric heavy chain.
 - [0130] In an analogous fashion, but using mRNA templates for cDNA construction for human kappa rather than γ chain, the expression plasmid for chimeric light chain is prepared.
- [0131] The foregoing two plasmids are then double transformed into <u>E. coli</u> W3110, the cells grown and the chains reconstituted as set forth in paragraph E.1-E.3 supra.

E.5 Preparation of Fab

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- E.5-1 Construction of a Plasmid Vector for Direct Expression of Murine Anti-CEA Gamma 1 Fab Fragment Gene pyCEAFabtrp207-1*
- [0132] Figure 13 presents the construction of p γ CEAFabtrp207-1*. 5 μ g of pBR322 was digested with Hind III, the cohesive ends made flush by treating with Klenow and dNTPs; digested with Pst I, and treated with BAP. The large vector fragment, fragment I, was recovered using 6 p rc nt PAGE followed by electro lution.
- [0133] 5 μg of pγCEAtrp207-1* was digest d with both BamH I and Pst I and the ~1570 bp DNA fragm nt (fragment II) containing the trp promot r and the gene sequence ncoding the variable region continuing into constant region and furth r into the anti-CEA gamma 1 chain hinge region, was isolated and purifi d after electrophoresis.
 - [0134] Expression of th anti-CEA gamma 1 chain Fab fragment rather than complet h avy chain r quires that a

termination codon be constructed at the appropriate location in the gene. For this, the 260 bp Nco I-Nde I DNA fragment from 20 μ g of the p γ 298 was isolated and purified. A 13 nucleotide DNA primer, the complement of which encodes the last 3 C-terminal amino acids of the Fab gene and 2 bases of the 3 needed for the stop codon, was synthesized by the phosphotries to remethod (supra). The probe hybridizes to nucleotides 754 to 767 (Figure 4) which has the following sequence:

AspCysGlyStop

5' GGGATTGTGGTTG 3'

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[0135] The third base of the stop codon is provided by the terminal nucleotide of the filled-in Hind III site from pBR322 cleavage described above. 500 ng of this primer was used in a primer repair reaction by phosphorylation at the 5' end in a reaction with 10 units T4 DNA kinase containing 0.5 mM ATP in 20 μ l, and mixing with ~200 ng of the Nco I-Nde I DNA fragment. The mixture was heat denatured for 3 minutes at 95° and quenched in dry-ice ethanol. The denatured DNA solution was made 60mM NaCl, 7mM MgCl₂, 7 mM Tris HCl (pH 7.4), 12 mM in each dNTP and 12 units DNA Polymerase I-Large Fragment was added. After 2 hours incubation at 37°C, this primer repair reaction was phenol, CHCl₃ extracted. ethanol precipitated, digested with BamH I and the reaction electrophoresed through a 6 percent polyacrylamide gel. ~50 ng of the 181 bp blunt end to BamH I DNA fragment, fragment III, was isolated and purified. [0136] ~100 ng of fragment I, ~100 ng each of fragments II and III were ligated overnight and transformed into E. coli K12 strain 294. Plasmid DNA from several tetracycline resistant transformants was analyzed for the proper construction and the nucleotide sequence through the repair blunt end filled-in Hind III junction was determined for verification of the TGA stop codon.

E.5.2 Production of Fab Protein

[0137] The plasmid prepared in E.5.1 is transformed into an <u>E. coli</u> strain previously transformed with pKCEAtrp207-1* as described above. The cells are grown, extracted for recombinant antibody chains and the Fab protein reconstituted as described in E.1.10.

Claims

 A non-glycosylated chimeric immunoglobulin species having specificity for a particular known antigen comprising chimericheavy and light polypeptide chains each having a constant region from a human antibody and a variable region from a murine antibody.

35 Patentansprüche

 Nichtglykosylierte, chimäre Immunglobulin-Spezies, die Spezifität für ein bestimmtes bekanntes Antigen aufweist, das chimäre schwere und leichte Polypeptidketten umfasst, die jeweils eine konstante Region von einem Human-Antikörper und eine variable Region von einem Mäuse-Antikörper aufweisen.

Revendications

 Espèce d'immunoglobuline chimérique non glycosylée ayant une spécificité pour un antigène connu particulier comprenant des chaînes polypeptidiques lourde et légère chimériques, chacune ayant une région constante d'un anticorps humain et une région variable d'un anticorps de murin.

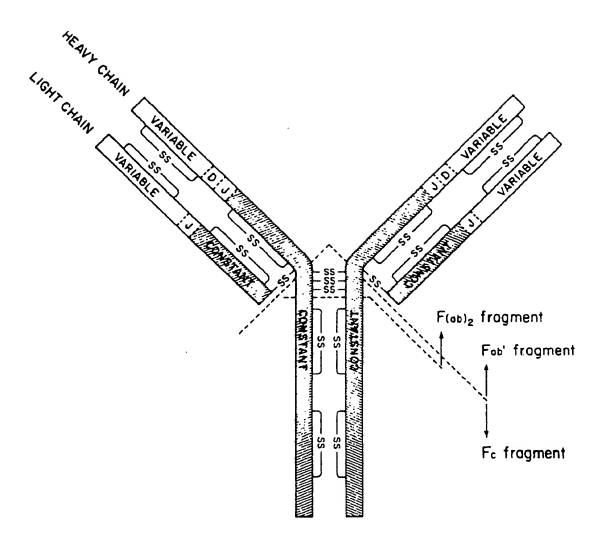


Fig. 1.

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501	hga I GCGTCAAA	sau3A dpn1 hgal GCGTCCTGAA CAGTTGGACT GATCAGGACI	JJA NI bby mall hincil Traggaca gcaaagacag cacctacag atgagcagca cctcaggt gaccaagga gagtatgaac gacayaacag	CACCTACAGO	fnu4HI bbv ATGAGCA	mnll hinc CCCTCACGTT	f 1 GACCAAGGAC	GAGTATGAAC	aluī Gacataacas	
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701	alui ali Agctcccag Tcgagggtc Foki	mnli mnli mnli hgta mboli ddel mnli mnli mnli hgta mnli mnli mnli hgta mnli mnli mnli mnli mnli foccoccag ciccatcota totoccto coccaccoccag ciccas ciccatcota totocctoc coccaccoccag coccaccoccag coccaccoccag foccaga da	ddel C taaggictig G attccagaac	mnll Gaggettee Cteegaagg	CACAAGCGAC GTGTTCGCTG	CTACCACTGT GATGGTGACA	hgia TGCGTGCTC ACGCCACGAG	mnli mnli caacctcct gttggagga	mnll CCCCACCTCC GGGGTGGAGG	
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-9 leu leu trp 1 UVG CVG UGG V	thr	ser thr arg his thr gly . UCC ACC CGG CAC ACU GGA	a s p G A C	a	pro lys asp fle asm val lys trp lys CCC AAA GAC AUC AAU GUC AAG UGG AAG	ser thr tyr ser met ser ser thr leu AGC ACC UAC AGC AUG AGC AGC ACC CUC	thr	GGUCUUGGAGGCUUCCCCACAAGCGACCUACCACI	UAUUCAAUAAAGUGAGUCUUUGCACUUGA
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sau96 hinfl avall mnll GAGTCAGCAC TGAACACGGA CCCCTCACGA TGAACTTCGG GCTCAGCTTG CTCAGTCGTG ACTTGTGCCT GGGGAGTGCT ACTTGAAGCC CGAGTCGAAC	AAACTCTCCT TTTGAGAGGA	GTAGTGGTGG CATCACCACC		mnll ddel: hphl CCTCA GTCACGTCT GCAGT CAGTGGCAGA
TGAACTTCGG ACTTGAAGCC	scrfl sau96 mull ecoRII avalI TGGAGCCTGG AGGTCCCTG ACCTCGGACC TCCCAGGGAC	GCAACCATTA	rsal Igtaccigc aaatgagcag Acatggacg titactcgtc	mn 1 1 dde 1 AGGAACCTCA ICCTTGGAGT
96 II mnli CCCCTCACGA GGGGAGTGCT		GGAGTGGGTC CCTCACCCAG	rsal Ctgtacctgc Gacatggacg	ACTGGGGTCA TGACCCCAGT
Sau96 avali TGAACACGGA CO ACTTGTGCCT GO	hinfi GGAGTCTGGG GGAGTCTTAA CCTCAGACCC CCTCAGAATT	i mnll mboll AGAAGAGGCT TCTTCTCCGA	GAGACAATGC CAAGAACACC CT CTCTGTTACG GTTCTTGTGG GA	GCTATGGACT CGATACCTGA
hinfl GAGTCAGCAC CTCAGTCGTG	hinfl GGAGTCTGGG CCTCAGACCC	hpali hinfl mt cagacteegg aga	GAGACAATGC CTCTGTTACG	AGCGGACTAT TCGCCTGATA
-	101	201	301	401

Fig. 4A.

			mn11 ra1 CG	
CTGTCCAGCG Gacaggtcec	hph! CCGTCACCTG GGCAGTGGAC	AGAAGTATCA TCTTCATAGT	sau3A dpn1 m foki ava1 CATCAGCAAG GATGATCCCG GTAGTCGTTC CTACTAGGGC	ddel Gytcaacagc actitccgci cagtcagtga Caagtigtgg Igaaaggcga gtcagtcact
xholl sau3A dpn1 bamh1 ctctggatcc	sau96 hae111 66 CCCAGCGAGA ICC GGGTCGCTCT	rsai Gtacagteee Catgteagg		dd actteceet Tgaargecoa
SCFI ECORII CCAGTGACAG TGACCTGGAA GGTCACTGTC ACTGGACCTT	sal mnll hat cagcccrces GTCGGAGCC	ndel TGGTTGTAAG CCTTGCATAT ACCAACATTC GGAACGTATA	accI TTGTGGTAGA AACACCATCT	GTTCAACAGC CAAGTTGTCG
scrFI ifaNI fokI if aNI scrFI il ecoRII scrFI eCORII ecoRII ECORII eCORII CACCTG GGATGCCTGG TCAAGGGCTA TITCCCTGAG CCAGTGACAG TGACCTGGAAA	finu4HI bbv ddel bbv ddel bbv ddel bbv ddel alul ctgrccacac ctrccact strctscast treactry cacretaac actrcacte caccette General Caccette General cacacacette caaggerea cacacgeta Gactgaaca treactry caccette	TGGTTGTAAG ACCAACATTC	hph! hinf! dde! fok! hg!A TCTTCCCCCC AAAGCCCAAG GATGTGCTCA CCATTACTCT GACTCTAAG GTCACGTGTG TIGTGGTAGA AGAAGGGGGG TTTCGGGTTC CTACACGAGT GGTAATGAGA CTGAGGATTC CAGTGCACACATCT	nn 1 I SAGGAGCA STCCTCGT
scrfi hi ecokii scrfi teii ecokie corfi teii ecorii scrfi Tgacctig ggatgctigg tcaagggta tttcctigag actgggac cctacggacc agttcccgat aaagggactc	fnu4H1 hbv dde1 a1u1 GC AGCTCAGTGA CG TCGAGTCACT	I fnu4HI bbv GCAGCACAA GGTGGACAAG AAAATTGTGC CCAGGGATTG CGTCGTGGTT CCACCTGTTC TTTTAACACG GGTCCCTAAC	mstII hinfl ddeI GACTCCTAAG	smal scrfi scrfi ncil ncil hpall Acccaccc GG(TGCGTGGG CC(
I TCAAGGGCTA AGTTCCCGAT	fn hb ddel Cactctgagc GTGAGACTCG	AAAATTGTGC TTTAACACG	I CCATTACTCT GGTAATGAGA	dde! alu! h cacactcag
foki 11 scrFI 6GAIGCCIGG	mnll CTGACCTCTA GACTGGAGAT	GGTGGACAAG CCACCTGTTC	hphl oki hgía gatgtgctca ctacacgagt	mnlI hgiA TGGAGGTGCA C ACCTCCAGG
scrfi sfaNi f hphi ecoRii bstEli GGTGACCTG G	PSt1 GTCCTGCAGT CAGGACGTCA	1 fnu4H1 bbv GCAGCACCAA CGTCGTGTT	AAAGCCCAAG TTTCGGGTTC	pvuli alui Cagctgatit Gtagatgatg Gtgaccaaa Catctactac
ncol CTAACTCCAT GATIGAGGTA	PVUII alui cttcccact gaaggtcga	scrFi haelli ncil hpall CACCCGCCA G	mboli Tcttccccc Agaagggg	pvuli alui cagetgattt GTGACCAAA
scrFI ncol sfaNI fokI fnu4HI bbv GCIGCCCAAA CTAACTCCAI GGGACCCTG GGATGC	hg1A GTGTGCACAC CACACGTGTG	SCFFI haeli ncfI bgll hpail CAACGTTGCC CACCCGGCCA	mboll r TCTGTCTTCA 1 AGACAGAAGT A	sauge avail AGGTCCAGTT TCCAGGTCAA
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CATCTCCAAA Gtagaggttt	ATAACAGACT Tattgtctga	GCTCTTACTT CGAGAATGAA	#b ddej CCATACTGAG GGTATGACTC	CTGTATAAT Gacatattta	
taqi Tcgagaaaa Agctctttt	GACCIGCATG	AACACGAATG TTGTGCTTAC	II TGCACAACCA ACGTGTTGGT	mnll mnll ACACCTAECT CCACCCTEC TGTGGATGGA GGTGGGGAGG	
fnu4HI bby TGGCTCAA TGGCAAGGAG TTCAAATGCA GGGTCAACAG TGCAGCTTC CCTGCCCCCA TCGAGAAAC CATCTCCAAA ACCGAGTT ACCGTTCCTC AAGTTTACGT CCCAGTTGTC ACGTCGAAAG GGACGGGGT AGCTCTTTG GTAGAGGTTT	haell! hae! cactcccaa ggaggagatg gccaaggata aagtcagtt gacctgcatg ataacagact gtggaggtt cctcgtctac ggitctat ttcagtcaga ctggacgtac tattgtctga	fnu4H1 bby TCTTCCCTGA AGACATTACT GTGGAGTGGC AGTGGAATGG GCAGCCAGCG GAGAACTACA AGAACATCA GCCCATCATG AACACGAATG GCTCTTACTT Agaaggaact tctgtaatga cacctcacgg tcaccttacc cgtcggtcgc ctcttgatgt tcttgtgagt cgggtagtac ttgtgcttac cgagaatgaa	sau96 mnll mnll ddel CGTCTACAGC AAGTCAATG IGCAGAAGAG CAACIGGGGG GCAGGAAATA CITICACCIG CICIGIGITA CATGAGGGCC IGCACAACCA CCATACIGAG GCAGAIGICG ITCGAGITAC ACGICIICIC GITGACCCIC CGCCIITAI GAAAGIGGAC GAGACACAAI GIACICGGG ACGIGIIGIG GGIAIGACIC		
fnu4HI bbv alui TGCAGCTTTC ACGTCGAAAG	haeili ali 6 GCCAAGGATA C CGGTTCCTAT	ddel Agaacactca TCTTGTGAGT	CTCTGTGTTA	hinfi CAGGAETCTG GTCCTGAGAC	
hinell GGGTCAACAG CCCAGTTGTC	hae hael ball GGGCAGATG G	GAGAACTACA	hph! CTTCACCTG GAAGTGGAC	sau96 11 avall 1CTGGTCTA	
TTCAAATGCA AAGTTTACGT	mn)I Cacctcccaa GTGGAGGGTT	fnudH1 bbv GCAGCCAGCG	I GCAGGAAATA CGTCCTTTAT	CTTGGAGCCC	
TGGCAAGGAG ACGTTCCTC	rsal ig tacaccattc ic atgiggtaag	AGTGGAATGG TCACCTTACC	mn ¹ CAACTGGGAG GTTGACCCTC	sau3A dpn1 GA TCCCAGIGTC CT AGGGTCACAG	
1 ACTGGCTCAA TGACCGAGTT	TCCACAGGTG AGGTGTCCAC	GTGGAGTGGC CACCTCACGG	mboll Tgcagaagag Acgtcttctc	scrfi ecorii da cc tegtaaatea eg accatttact	GGGAAAAA
SCPFI CORRII ACTTCCCATC ATGCACCAGG ACTGGCTCAA TGAAGGGTAG YACGTGGTC TGACCGAGTT	rsal Accaaaggca gaccgaaggc Tccacaggtg Tacaccattc Cacctccaa Tgrittcgt Ctggcttcg Aggtgtcac Atgtggtaag Gtggaggtt	oli Agacattact Tctgtaatga	alui Aagctcaatg Ttcgagttac	sc ec cccactctcc gggtgagg	AAAGCACCCA GCACTGCCTT GGGAAAA TITCGTGGGT CGTGACGGAA CCCTTTTT
ACTTCCCATC TGAAGGETAG	ACCAAAGGCA TGGTTTCCGT	mboli TCTTCCCTGA AGAAGGGACT	acci CGTCTACAGC GCAGATGTCG	mnli Aagagcetet TTCTCGGAGA	AAAGCACCCA TTTCGTGGGT
1001	1101	1201	1301	1401	1 50 1

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glu GAA	AGA	S SET	D thr	o	s leu c cug	0 u 91n G CAG	0 & Ja 6 GCC	o Pre C uuc
cy s ugu	30 ser AGU	60 CCA	90 8 8 P	120 thr	150 Cys	180 1 eu	210	240 7 (8)
gla CAG	phe	Jeu	9) u GAG	val GUC	9 1 y	va) GUC	b1s CAC	ser
va) 600	thr	h 1 s C A C	ser ucu				8 1 8 GCC	ser uca
val GUU	phe UUC	Ser	a rg AGG	thr	thr	0 7 0 C C A	, e] GUU	va] GUA
1ys AAA	9	ser AGU	Jeu CUG	9 1 y 6 G A	v a l GUG	phe uuc	AAC	
leu UUA	ser ucu	9 y 6 G U	ser AGU	gln CAA	met AUG	t hr	0 y S UGC	Pro
val GUU	a) a	91y GGU	ser AGC	9 1 y GGU	ser UCC	h i s CAC	thr	**) 600
leu CUG	ala ala GCA GCC	ser AGU	ae t AUG	trp UGG	asn AAC	√a } 6116	va] GUC	ACA
val GUC	cys UGU	ser Agu	9 n CAA	ty r UAC	140 1 ala gin thr a 1 GCC CAA ACU A	9	th And	cys UGU
-10 1eu	ser UCC	11e AUU	leu CUG	a s p G A C	9 1 n C A A	Ser	9 1 u GAG	11e AUA
tyr UAC	20 leu CUC	50 thr Acc	80 tyr UAC	110 metasp AUG GAC	140 ala 600	170 ser ucc	200 ser AGC	230 cys UGC
1) e AUU	1ys Aaa	8 1 8 GCA	Jeu CUG	a 1 a 5 C U	1 S	leu CUG	979 CCC	022
leu vug	Je v CUG	∨a	thr		ser ucu	Ser	87.8 000	lys pro
Ser	ser	trp UGG	AAC	asp GAC	91y 66A	74	53	cy s ugu
leu CUC	9 J 7 S	9 1 u G A G	ala lys GCC AAG	val ala d GUA GCG	ala pro GCC CCU	in ser g	ser pro	9 1 y 66 U
91y 666	9 J y GGA	leu cu6	8 0 0 0	val GUA	ala 339	20	Ser	o y s
phe	010 CCU	4 A	asn AAU	Jeu UUA	le v CUG	trp UGG	P 7 0	ASPGAU
asn	91 u GA G	1ys AAG	a s b	ser ucg	Pro	thr	val GUC	8 8 9 9 9 9
met AUG	met Aug	9 J u G A G	a rg AGA	17e AUU	tyr UAU	va! GUG	thr ACU	010
ACG	leu UUA	9 T 0	ser UCC	leu Cưu	va] GUC	thr	va) 606	v a ₹ G U G
ວກວວ	10 val GUC	40 thr ACU	70 11e AUC	100 Pro CCU	130 ser ucu	160 val GUG	190 ser uca	220 11e AUU
GACC	9] y GGA	gln CAG	thr	979 000	Pro CCA	pro	s e r A G C	1ys AAA
CACG	91y 666	a r g CGC	phe UUC	a rg AGA	919 555	9 1 c	Ser	lys AAG
UGAA	ser ucu	va) GUU	a rg CGA	a 1 a G C A	thr	9 ro	Jeu CUG	a sp GAC
GCAC	glu GAG	trp UGG	9] y 666	cys ugu	thr ACG	p he UUC	thr ACU	va] GUG
GAGUCAGCACUGAACACGGACCCCUCACG	va] 606	ser UCU	9	tyr UAC	1 y s	tyr	tyr UAC	1ys AAG
G.A.	Jew	me t AUG	cys 060	tyr UAU	a] a 6C C	9 J y 66 C	Jeu CUC	thr
	AUG	8 1 8 6 C C	gla CAG	met AUG	Ser	1 y s A A G	a sp GAC	Ser
	505	tyr	ACA	a] a	ser ucc	va) GUC	ser ucu	Ser

Fig. SA

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0 43		0 0 0 0 0 0 0 0	O S a S D G GAU			UCCCAGUGUCCU	
270 asp	300 1 Phe	-	360 ala lys GCC AAG	390 858	420 91y 66A	5V	
asp	thr	010	8 J 8	9 1 u 6 A G	8 1 8 1 8 1 8 1 8 1 8 1 8 1 8 1 8 1 8 1		
1ys AAG	AGC		met AUG	8 1 8 6 7 6	91 u	6 2 6 2	
Ser	AAC	000 e	g i n CAG	P 70	try UGG	447 145 AAA	
asp fle ser lys GAC AUC AGC AAG	or co	ralaala Jecaecu	9 1 e GA 6	91 n	AAC	9 5 5 5 5	
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4.8.1 6.0.A	9	g val asm ser G guc aac agu	979	AAU	1ys AAG	ser	
va 1 606	9 } c	va] GUC	Pro CCU	200	91n CAG	A T S	¥ ¥
val val val a	arg glu glu gln phe asn CGG GAG GAG CAG UUC AAC	310 his gin asp trp leu asn gly lys glu phe lys cys arg CAC CAG GAC UGG CUC AAU GGC AAG GAG UUC AAA UGC AGG	340 1ys gly arg pro lys ala pro gln val tyr thr ile pro pro pro lys glu gln AAA GGC AGA CCG AAG GCU CCA CAG GUG UAC ACC AUU CCA CCU CCC AAG GAG CAG	370 thr asp phe pro glu asp fle thr val glu trp gln trp asn gly gln pro ACA GAC UUC UUC CCU GAA GAC AUU ACU GUG GAG UGG CAG UGG AAU GGG CAG CCA	400 thr asm gly ser tyr phe val tyr ser lys leu asm val glm lys ser asm trp ACG AAU GGC UCU UAC UUC GUC UAC AGC CUC AAU GUG CAG AAG AGC AAC UGG	430 glu gly leu his asm his his thr glu lys ser leu ser his ser pro gly lys e GAG GGC CUG CAC AAC CAC CAU ACU GAG AAG AGC CUC UCC CAC UCU CCU GGU AAA	GAAA
250 val leu thr 11e thr 1eu thr pro lys val thr cys ' GUG CUC ACC AUV ACU CUG ACU CCU AAG GUC ACG UGU	290 gin pro 6	Cy 5 UGC	1) e AUU	4 T D	AAU	le CUC	UUGG
260 thr ACG	290 91n CAA	320 1 y s AAA	350 thr ACC	380 914 6AG	410 Jeu CUC	440 ser AGC	3390:
val 600	thr	e y o	tyr UAC	va 1 60 G	1ys AAG	1ys AAG	6CAC
1ys AAG	6 P P P P P P P P P P P P P P P P P P P	97 u	val GUG	thr	Ser	g] u GAG	CCCA
Pro	a]a 600	1ys AAG	(A6	11 e Auŭ	tyr UAC	thr	AGCA
thr	thr ala gin thr ACA GCU CAG ACG	9 } y	Pro	a s b GAC	va] GUC	h is	AUAA
1eu CUG	VOT NES	asa AAU	8)8 GCU	69 - CA B	phe UUC	h1s CAC	AUAA
thr	606	leu CUC	1ys AAG	010	tyr UAC	ASB	กอกว
11e AUU	280 asp asp val glu v GAU GAU GUG GAG E	trp UGG	970	Phe	ser	h s CAC	ວວຄວ
thr	606	a s p G A C	a rg Aga	phe CUC	9 5 66 0	leu CUG	ACCC
Jeu	a S P G A U	gla	91x 66c	8 8 0 0 A 0	AAU	91y 66c	ວວກວ
250 val GUG	280 asp GAU	310 h1s cac	340 7ys AAA	370 thr ACA	400 thr ACG	430 9)u 6AG	CUAC
asp GAU	val GUA	met AUG	thr	11e AUA	AAC	h i s CAU	ACAC
lys AAG	p he UUU	11e AUC	1 y S A A A	met Aug	ae t AUG	Jeu	ეიიც
070	trp UGG	pro 11e CCC AUC	ser	cy s UGC	11e AUC	ser val ucu gug	GGAC
1ys AAG	Ser	Jeu	11e AUC	thr	000	ser	UACA
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D he UUC	7 8 7 0 8 C	789	9 n c g	va) 60C	AAC	oppe (13335
1) e	0 4 5 2 4 5 5 4 5 5 4 5 5 5 5 5 5 5 5 5 5	Ser	AUC	AAA	AAG 1	thr Acu (UGGAGCCCUCUGGUCCUACAGGACUCUGACACCUACCUCCACCCCUCCCU
		_	-	_			_

Fig. 5B.

